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Guo, Qiang; Meng, Lin; Humphreys, Mike W.; Scullion, John; Mur, Luis A.J.

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1 **Title:** Expression of *FIHMA3*, a vacuolar P_{1B2}-ATPase from *Festulolium loliaceum*,
2 correlates with response to cadmium stress

3 **Authors:** Qiang Guo^a, Lin Meng^{a*}, Mike W. Humphreys^b, John Scullion^b, Luis A. J.
4 Mur^b

5 **Institutional addresses:** ^a Beijing Research and Development Center for Grass and
6 Environment, Beijing Academy of Agriculture and Forestry Science, Beijing 100097, P.
7 R. China; ^b Institute of Biological, Environmental and Rural Sciences, Aberystwyth
8 University, Aberystwyth SY23 3DA, UK

9 ***Corresponding author:** Lin Meng; E-mail: menglin9599@sina.com

10 **Tel:** +86-10-51503345

11 **Fax:** +86-10-51503297

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Abstract

Heavy metal ATPase 3 (HMA3), a P_{1B2}-ATPase, is a key tonoplast transporter involved in mediating the vacuolar sequestration of cadmium (Cd) to detoxify the intake of this element by plants. HMA3 expression in response to Cd stress has not been previously examined in the grass hybrid species *Festulolium loliaceum* (Huds.) P. Fourn. In this study, *FlHMA3* isolated from *F. loliaceum* was found to comprise 833 amino acid residues with 77% homology to the rice *OsHMA3*. Transient expression of *FlHMA3* fused to enhanced green fluorescent protein in *Arabidopsis* protoplasts suggested its localization to vacuolar membranes. Quantitative real-time RT-PCR analysis of *F. loliaceum* revealed that *FlHMA3* is expressed predominantly within roots and up-regulated by excess Cd. Over the 168 h treatment, Cd content of *F. loliaceum* roots was significantly higher than that of shoots, regardless of external CdCl₂ concentrations. A significant positive correlation was found between *FlHMA3* expression and Cd accumulation in roots of *F. loliaceum* seedlings subjected to 10–100 mg L⁻¹ CdCl₂ for 168 h or, in a separate experiment, to 25 or 100 mg L⁻¹ CdCl₂ for the same duration. These findings provide evidence that *FlHMA3* encodes a vacuolar P_{1B2}-ATPase that may play an important role in Cd²⁺ sequestration into root cell vacuoles, thereby limiting the entry of Cd²⁺ into the cytoplasm and reducing Cd²⁺ toxicity.

Keywords: cadmium, *Festulolium loliaceum*, *FlHMA3*, phytomediation, vacuole sequestration

1. Introduction

An expanding global population and the limited availability of agriculturally suitable land have stimulated interest in the agricultural potential of more marginal areas and contaminated “brown-field” locations on former industrial sites. Of the estimated 30,000 t of Cd released annually into the environment, 13,000 t are due to human activity (Gallego et al., 2012). A non-essential heavy metal with high toxicity to plants, Cd interferes with the homeostasis of essential elements such as zinc, calcium, and iron and initiates their displacement from proteins (Verbruggen et al., 2009; Guo et al., 2014). Cd contamination of soil has become a serious environmental concern and also threatens human health via its accumulation in the food chain (Satoh-Nagasawa et al., 2012). Crops and products derived from livestock raised on plant-based diets are important sources of heavy metals absorbed by humans (Peralta-Videa et al., 2009). Understanding the mechanism of Cd accumulation in plants and the factors affecting its deposition are crucial to reduce entry of Cd into the human food chain.

Plants have evolved alternative adaptation strategies to cope with Cd stress. One such mechanism involves production of phytochelatins (PCs), which are glutathione-derived peptides. PCs are synthesized in the cytosol, where they form PC-Cd complexes that are subsequently sequestered into vacuoles to reduce the deleterious effects of Cd accumulation in the cytosol (Mendoza-Cózatl et al., 2005; Clemens, 2006). This process plays an important role in heavy metal homeostasis and detoxification (Colangelo and Guerinot, 2006; Hanikenne and Nouet, 2011). Generally, vacuolar sequestration of Cd^{2+} can be mediated by heavy metal

1 transporters in plants (Hirschi et al., 2000; Song et al., 2003; Korenkov et al., 2007;
 2 Wojas et al., 2009). Among these transporters are P_{1B}-ATPases (heavy metal
 3 P_{1B}-ATPases, HMAs), a large group of ATP-driven pumps implicated in the transport
 4 of monovalent Cu⁺/Ag⁺ (P_{1B1}-ATPases) and divalent Zn²⁺/Cd²⁺/Co²⁺/Pb²⁺
 5 (P_{1B2}-ATPases) heavy metal cations across plant membranes (Williams and Mills,
 6 2005). As is well known, HMAs are involved in removal of heavy metal ions from the
 7 cytosol into either the apoplast, the vacuole, or into other organelles (Hussain et al.,
 8 2004; Andrés-Colás et al., 2006; Kim et al., 2009). In *Arabidopsis thaliana*,
 9 *AtHMA1-AtHMA4* and *AtHMA5-AtHMA8* transport divalent and monovalent cations,
 10 respectively (Cobbett et al., 2003; Williams and Mills, 2005). *AtHMA3* belongs to the
 11 Zn²⁺/Cd²⁺/Co²⁺/Pb²⁺ subgroup; localized in the tonoplast, it helps detoxify essential
 12 biological (Zn²⁺) and non-essential (Cd²⁺, Co²⁺, and Pb²⁺) heavy metals by
 13 participating in their vacuolar sequestration (Morel et al., 2009). *AtHMA3* is a major
 14 locus in *A. thaliana* responsible for the regulation of Cd accumulation (Chao et al.,
 15 2012). *AhHMA3* has high-level constitutive expression in *A. halleri*, a Zn
 16 hyperaccumulator and relative of *A. thaliana*, which suggests that this gene is
 17 involved in high Zn accumulation (Becher et al., 2004). Similarly, *TcHMA3* from
 18 *Thlaspi caerulescens*, a Cd hyperaccumulator, is highly expressed in leaves; it plays
 19 an important role in the detoxification of Cd by sequestering Cd into leaf vacuoles,
 20 thereby contributing to Cd hyperaccumulation and hypertolerance (Ueno et al., 2011).
 21 Among the nine *HMA* genes identified in rice is *OsHMA3*, a member of the
 22 Zn²⁺/Cd²⁺/Co²⁺/Pb²⁺ subgroup (Miyadate et al., 2011; Takahashi et al., 2012).

1 *OsHMA3*, a tonoplast-localized transporter of Cd within root cells, plays a role in the
 2 sequestration of Cd²⁺ into root cell vacuoles (Ueno et al., 2010). *OsHMA3* has been
 3 identified as the locus responsible for regulation of Cd accumulation in shoots of rice
 4 cultivars Anjana Dhan and Cho-Ko-Koku. When the function of this protein is lost,
 5 Cd²⁺ passage through the xylem is increased, thereby leading to Cd²⁺ accumulation in
 6 the shoots (Ueno et al., 2010; Miyadate et al., 2011). Sasaki et al. (2014) recently
 7 found that overexpression of *OsHMA3* contributed to reduced Cd²⁺ accumulation in
 8 the grain and to enhanced Cd tolerance in rice. Taken together, the available evidence
 9 implies that *HMA3*-mediated vacuolar sequestration of heavy metals plays an
 10 important role in metal detoxification.

11 *Festulolium* grass hybrids combine many of the attributes of *Lolium* species
 12 (ryegrasses), such as high growth rates that provides high yields of nutritious,
 13 palatable fodder for livestock, and those of *Festuca* species (fescues) which provide
 14 resilience against different climatic and edaphic stresses (Humphreys et al., 2014).
 15 Natural *Festulolium* species hybrids exist, especially in undisturbed marginal
 16 grassland locations frequently exposed to stress conditions where productive *Lolium*
 17 species would be more disadvantageous (Humphreys et al., 1995). A range of
 18 synthetic *Festulolium* species hybrids are also being generated by grass breeders to
 19 achieve productive, stress-adapted varieties suitable for agriculture, with safeguards to
 20 assist withstanding various climatic conditions. The natural grass species hybrid
 21 *Festulolium loliaceum* survives in waterlogged soils prone to flooding and its
 22 synthetic varieties developed combat freezing temperatures or to mitigate incidents of

1 flooding (Macleod et al., 2013). *Festulolium loliaceum* was developed following the
2 hybridisation of *Lolium perenne* (perennial ryegrass) with *Festuca pratensis* (meadow
3 fescue). Eventhough *F. loliaceum* has been broadly recognized for its resilience to
4 stress conditions, studies in this regard have been primarily confined to climatic rather
5 than edaphic stresses; consequently, the potential use of this species for
6 bioremediation and its tolerance to Cd-contaminated soils have not been previously
7 explored. In this context, we isolated and characterized *FIHMA3* from *F. loliaceum*
8 and verified its subcellular localization. We also analyzed *FIHMA3* expression in
9 plants exposed to high Cd concentrations, which revealed patterns consistent with a
10 role in the conferral of heavy metal tolerance. Our findings provide useful initial
11 information on the potential future agricultural application of this grass hybrid on
12 Cd-contaminated soils.

2. Materials and Methods

2.1 Plant growth conditions and treatments

Seeds of *Festulolium loliaceum* variety Prior ($2n = 4x = 28$) were provided by the Institute of Biological, Environmental, and Rural Sciences (IBERS), Aberystwyth University. Seeds were sterilized with 5% sodium hypochlorite solution for 5 min, rinsed thoroughly with distilled water, and then germinated on moistened filter paper for 168 h at 25°C in dark. After emergence of plumules, uniform seedlings were selected and transferred into plastic containers filled with 0.6 L modified Hoagland's solution containing 2 mM KNO₃, 1 mM NH₄H₂PO₄, 0.5 mM Ca(NO₃)₂·4H₂O, 0.5 mM MgSO₄·7H₂O, 60 µM Fe-citrate, 92 µM H₃BO₃, 18 µM MnCl₂·4H₂O, 1.6 µM ZnSO₄·7H₂O, 0.6 µM CuSO₄·5H₂O, and 0.7 µM (NH₄)₆Mo₇O₂₄·4H₂O for 5 weeks. The nutrient solution was renewed every 2 d. All seedlings were grown in a CE growth chamber under a 16 h/8 h day/night cycle at 25°C/18°C, a relative humidity of 50% to 60%, and a light intensity of 200 µmol m⁻² s⁻¹. Five-week-old plants were used for all treatments, with each treatment replicated eight times and each replicate comprising five individual plants. Containers for all treatments were arranged in a completely randomized block design. Two treatment approaches were used: (i) Hoagland's nutrient solution supplemented with 0, 10, 25, 50, and 100 mg L⁻¹ CdCl₂ for 168 h; and (ii) Hoagland's nutrient solution containing 25 or 100 mg L⁻¹ CdCl₂, with plants removed following 0, 3, 6, 12, 24, 48, 72, 96, 120, 144, and 168 h exposure. These exposure concentrations were not selected to simulate field conditions. Rather they were determined to observe Cd uptake and stress of the plants over a short period of exposure in order to better understand plant mitigation

1 responses.

2 2.2. Calculation of relative growth rate

3 The relative growth rate (RGR) of whole plants was calculated using the formula
4 $RGR = (\ln W_j - \ln W_i) / \Delta t$, where W_i and W_j are dry weights before and after 168 h
5 treatment, respectively, and Δt is elapsed time between the two measurements
6 (Martínez et al., 2005).

7 2.3. Cloning of *FlHMA3*

8 Total RNA was extracted according to Guo et al. (2012) from roots of *F.*
9 *loliaceum* seedlings exposed to 100 mg L⁻¹ CdCl₂ for 24 h. First-strand cDNA was
10 synthesized from 2 µg total RNA using an oligo(dT) primer and PrimeScriptRTase
11 (Takara). The partial cDNA fragment of *FlHMA3* was amplified by PCR using
12 degenerate primers P1 and P2 (Table S1) designed based on the gene sequences of
13 *BdHMA3* (*Brachypodium distachyon*, XM_003561234), *HvHMA3* (*Hordeum vulgare*,
14 KU212808), *OsHMA3* (*Oryza sativa*, XM_015791882), *TaHMA3* (*Triticum aestivum*,
15 KF683298), and *ZmHMA3* (*Zea mays*, XM_008671782). PCR cycling conditions
16 were as follows: 94°C for 2 min, followed by 30 cycles of 94°C for 30 s, 55°C for 30
17 s, and 72°C for 40 s, with a final extension at 72°C for 10 min. The PCR product was
18 purified from agarose gels, ligated into a pMD-19T vector (Takara), and sequenced by
19 Sangon Biotech (China). The 5'- and 3'-ends of *FlHMA3* were obtained with 5'- and
20 3'- Rapid Amplification of cDNA Ends kits (SMARTer RACE, Clontech) according
21 to the manufacturer's instructions and specific primers P3 and P4, respectively (Table
22 S1). These fragments were assembled to obtain the full-sequence of *FlHMA3* cDNA.

2.4. Sequence analysis

A BLAST search was performed using the NCBI platform (<http://www.ncbi.nlm.nih.gov/BLAST>). Sequence analysis of cDNA and multiple alignments were performed with DNAMAN 8.0 software. Molecular mass and isoelectric point of the deduced protein encoded by *FlHMA3* was predicted using the ExPASy proteomics server (<http://www.expasy.org>). Hydrophobicity values were calculated using the program TMPRED available at <http://www.ch.embnet.org/>. Phylogenetic relationship of *FlHMA3* with other plant *HMA*s multiple sequence alignment was analyzed by multiple alignments using Clustal X software (Thompson et al., 1997). Then a phylogenetic tree was constructed by MEGA6.0 software using the neighbor-joining method with 1,000 bootstrap replicates (Tamura et al., 2011).

2.5. Subcellular localization of *FlHMA3*

The open reading frame (ORF) of *FlHMA3* excluding the stop codon was amplified using PrimeSTAR HS DNA polymerase with primers P5 (Table S1, EcoRI restriction site underlined) and P6 (Table S1, KpnI restriction site underlined), cloned into a pMD-19T vector, and sequenced by Sangon Biotech. The amplified fragment was cut from the pMD-19T plasmid using EcoRI and KpnI restriction enzymes and cloned into a pBSHES-NL vector to generate a fusion with enhanced green fluorescent protein (EGFP) under the control of the CaMV35S promoter. The *FlHMA3*-GFP fusion construct or a non-GFP-tagged vector construct was transiently expressed in protoplasts isolated from *A. thaliana* Col-0 cell suspensions using the polyethylene glycol-mediated method (Yoo et al., 2007). Protoplasts containing the

1 plasmids were incubated at 23°C for 2–3 d in darkness. For FM4-64 staining,
 2 protoplasts were transferred into 50 μ M FM4-64 in Murashige-Skoog medium
 3 containing 0.4 M mannitol for 10 min at 4°C according to the method of Ueda et al.
 4 (2001). Fluorescent signals from both GFP and FM4-64 in the protoplasts were then
 5 observed using an inverted Carl Zeiss LSM 710 confocal laser scanning microscope.
 6 GFP and FM4-64 were excited at 488 nm and 543 nm, respectively, with their
 7 corresponding fluorescence emission signals detected between 498–539 nm and
 8 580–650 nm, respectively.

9 2.6. Expression analysis of *FlHMA3*

10 Total RNA was extracted with Trizol kit (Takara) following the manufacturer's
 11 instructions. First-strand cDNA was synthesized from 2 μ g of total RNA using an
 12 oligo(dT) primer and PrimeScriptRTase (Takara). Quantitative real-time RT-PCR
 13 (qRT-PCR) was performed using SYBR Premix Ex *Taq* II (Perfect Real Time) (Takara)
 14 on a StepOnePlus Real-Time PCR system (ABI) to monitor the amplification of each
 15 cDNA fragment. qRT-PCR amplification of *FlHMA3* was carried out with the primer
 16 pair P7 and P8 (Table S1), which yielded a 225 bp product. For use as a reference in
 17 the qRT-PCR, a 130 bp region of the *actin* gene was amplified using primers A1 and
 18 A2, which were designed according to the partial cDNA sequence of *actin* from *F.*
 19 *lioliceum* (Table S1). Primer sequences were designed with Primer 5.0 software. The
 20 amplification protocol consisted of an initial denaturation step of 95°C for 10 min,
 21 followed by 40 cycles of 95°C for 30 s, 60°C for 30 s, and 72°C for 30 s. Gene
 22 expression data were normalized and relative expression was calculated by the $2^{-\Delta\Delta C_t}$

1 method (Livak and Schmittgen, 2001). Each experiment included three biological
2 replicates.

3 *2.7. Determination of Cd content*

4 Harvested plants were thoroughly washed with deionized water, divided into
5 shoots and roots, and oven dried at 80°C to a constant weight. Dried shoots and roots
6 were ground and digested in a mixture of HNO₃/HClO₄ (5/1, v/v) for 24 h and then
7 heated at 150–200°C to near dryness. After cooling, the residue was dissolved in
8 distilled deionized water to a total volume of 20 mL. Cd content was determined
9 using an atomic absorption spectrophotometer (AA-6300C, Shimadzu, Kyoto, Japan).

10 *2.8. Statistical analysis*

11 All data were calculated as means plus standard deviation (SD). Statistical
12 analyses, one-way analysis of variance, and Duncan's multiple range tests comparing
13 treatment means were performed using SPSSv13.0 statistical software (SPSS Inc.,
14 Chicago, IL, USA).

3. Results

3.1. Characterization of *FlHMA3*

The full-length cDNA of *FlHMA3* isolated from roots was obtained by RT-PCR and rapid amplification of cDNA ends. The sequence, which comprised 2,957 bp, contained a 2,502 bp ORF encoding 833 amino acid residues with an estimated molecular mass of 87.58 kDa and a theoretical isoelectric point of 5.33. As shown in Fig. 1, multiple sequence alignment revealed that *FlHMA3* shares high homology with amino acid sequences of orthologs *TaHMA3* from wheat (83%) and *OsHMA3* from rice (77%) and, to a lesser extent, *AtHMA3* from *A. thaliana* (54%). This similarity indicates that *FlHMA3* is a P_{1B2}-ATPase. A previous study has shown that *OsHMA3* encodes a divalent metal ion (Cd²⁺) transporter P_{1B2}-ATPase involved in mediating the sequestration of Cd into vacuoles (Miyadate et al., 2011). Our phylogenetic analysis placed *FlHMA3* into a clade with other *HMA3* genes from closely related monocotyledonous species such as wheat (*TaHMA3*) and rice (*OsHMA3*) (Fig. S1). These results suggest that *FlHMA3* encodes a P_{1B2}-ATPase transporter.

Hydrophobicity plot analysis of the deduced polypeptide showed that *FlHMA3* has eight transmembrane domains (Fig. 1, TM1 to TM8). Like all P-type ATPases, it contains the characteristic motifs for an ATP binding (GDGxNDx), a phosphorylation (DKTGTLT), and an HP locus forming a large cytoplasmic loop. Moreover, *FlHMA3* possesses a CPC ion transduction motif in TM5, and a heavy-metal-associated domain containing the motif GxCCxxE was located at the N terminus of *FlHMA3*.

3.2. Transient expression of *FlHMA3* in mesophyll protoplasts of *Arabidopsis*

To determine the subcellular localization of *FIHMA3*, a GFP reporter construct was developed to express a fusion protein consisting of GFP fused to the C terminus of *FIHMA3* (FIHMA3:GFP) under the control of the CaMV35S promoter. This construct was then transfected into mesophyll cells of *A. thaliana* protoplasts using the polyethylene glycol method. Expression of the control (35S:GFP) resulted in distribution of GFP signals throughout the cytoplasm and the nucleus of mesophyll protoplasts (Fig. 2A). When FIHMA3:GFP was transfected into the protoplasts of mesophyll cells, however, the GFP signals were clearly separated, with FM 4-64 signals in the area of the plasma membrane and tonoplast (Fig. 2B). This observation suggests that *FIHMA3*, in line with its presumed function, is localized specifically on the vacuolar membrane.

3.3. Expression patterns of *FIHMA3* in *F. liliaceum* exposed to Cd

To investigate the effect of Cd on transcription of *FIHMA3*, plants were treated with different concentrations of CdCl₂ for 168 h. As shown in Fig. 3A, mRNA levels of *FIHMA3* were 2.2–6.6 folds higher in roots than in shoots at external CdCl₂ concentrations of 10–100 mg L⁻¹. Compared with the control, exposure to 10–100 mg L⁻¹ CdCl₂ led to significantly increased *FIHMA3* transcription in roots, with the highest transcription levels observed with the most concentrated solution (100 mg L⁻¹ CdCl₂). The expression of *FIHMA3* was significantly lower in shoots and did not change in response to the increase in CdCl₂ concentration.

To further determine the kinetics for Cd²⁺ induced activation of *FIHMA3* in tissues, *FIHMA3* transcript levels were examined in plants exposed to 25 and 100 mg

1 L^{-1} CdCl_2 over a 168 h period. When exposed to 25 mg L^{-1} CdCl_2 , *FlHMA3* transcript
 2 levels increased gradually in both roots and shoots from 3 to 168 h, but the magnitude
 3 was higher in roots than in shoots (Fig. 3B). In the case of 100 mg L^{-1} CdCl_2
 4 treatment, *FlHMA3* transcript levels increased rapidly in both shoots and roots from 3
 5 to 72 h; after peaking at 96 h, levels declined and remained constant following
 6 exposure for 120–168 h. Under this treatment, *FlHMA3* expression was always higher
 7 in roots than in shoots (Fig. 3C). Taken together, these results provide strong evidence
 8 to suggest that *FlHMA3* at least within the 168 h timeframe used in this study is
 9 expressed primarily in roots, with the extent of its transcription determined by the
 10 level of CdCl_2 exposure.

11 3.4. Cd accumulation in tissues and its relationship with the expression levels of 12 *FlHMA3*

13 As external CdCl_2 concentrations were increased, Cd contents of shoots and roots
 14 increased progressively after 168 h of growth; however, Cd contents of roots were
 15 significantly higher than those of shoots at concentrations of $10\text{--}100 \text{ mg L}^{-1}$ CdCl_2
 16 (Fig. 4A). Moreover, an increase in CdCl_2 concentration from 50 to 100 mg L^{-1} had
 17 no impact on Cd contents of shoots (Fig. 4A). We further observed a significant
 18 positive correlation between Cd content and *FlHMA3* expression level in roots of *F.*
 19 *loliaceum* exposed to $25\text{--}100 \text{ mg L}^{-1}$ CdCl_2 for 168 h (Fig. 4B). Despite the increased
 20 *FlHMA3* expression, however, the increase in Cd had a negative impact on plant
 21 growth rate even though roots did not appear to be necrotic or changed in overall
 22 length or branching and were therefore considered to be functional. A 14%–38%

1 reduction in relative growth rate compared with the control was recorded between the
2 treatment extremes (10–100 mg L⁻¹ CdCl₂) (Fig. S2). Despite this effect the *F.*
3 *liiaceum* plants survived following the treatments without further evidence of
4 toxicity symptoms such as chlorosis or necrosis, consistent with our presumption that
5 root functionality was maintained.

6 To monitor the differences in Cd accumulation in root and shoot tissue over time,
7 we recorded changes in plants exposed to 25 or 100 mg L⁻¹ CdCl₂ over a 168 h period.
8 The addition of either 25 or 100 mg L⁻¹ CdCl₂ significantly increased Cd content in
9 both shoots and roots, but the content was always higher in roots than in shoots (Fig.
10 5A). In plants undergone 100 mg L⁻¹ CdCl₂ treatment, the concentration of Cd in both
11 shoots and roots was significantly greater than in tissues exposed to 25 mg L⁻¹ CdCl₂
12 from 3 to 168 h (Fig. 6A). In addition, we observed no significant difference in Cd
13 accumulation by shoots in the presence of either 25 or 100 mg L⁻¹ CdCl₂ between 48
14 and 168 h (Figs. 5A–6A). A significant positive correlation was found between Cd
15 content and expression levels of *FIHMA3* in roots of *F. loliaceum* exposed to either 25
16 or 100 mg L⁻¹ CdCl₂ concentrations within the 3–168 h time frame (Figs. 5B–6B).
17 This result provides supporting evidence to suggest an association between Cd
18 tolerance and *FIHMA3* expression in *F. loliaceum*.

4. Discussion

P_{1B}-ATPases are involved in heavy metal transport through biological membranes via an ATP-dependent process. P_{1B2}-ATPases, which are unique to plants and have attracted much attention, play a critical role in controlling the translocation of Zn²⁺ or Cd²⁺ from roots to shoots and in sequestration of Cd²⁺ from the cytoplasm into the vacuole (Cobbett et al., 2003; Morel et al., 2009; Mendoza-Cózatl et al., 2011). However, studies of similar proteins in non-model plants have been lacking. In this paper, we have presented the first characterization of a homologous P_{1B2}-ATPase from a synthetic hybrid of the forage grass species *F. loliaceum*, an amphiploid species hybrid of the agricultural grasses *Lolium perenne* and *Festuca pratensis*. FIHMA3 has amino acid homologies of 54% and 77%, respectively, with AtHMA3 of *A. thaliana* and OsHMA3 of rice, both belonging to Zn²⁺/Cd²⁺/Co²⁺/Pb²⁺ transporting group of P_{1B2}-ATPases (Gravot et al., 2004; Miyadate et al., 2011). The FIHMA3 polypeptide sequence was found to possess the expected features of eight transmembrane domains and a CPC motif. The motifs DKTGTLT, HP, and GDGxNDx shown in Fig. 1 are considered to be the domains of ion transduction, phosphorylation, and translocation of metal ions in the large cytoplasmic loop and for ATP binding. Similar motifs have been reported for AtHMA3 from *A. thaliana* (Gravot et al., 2004), GmHMA8 from soybean (Bernal et al., 2007), OsHMA3 from rice (Miyadate et al., 2011), TcHMA3 from *T. caerulescens* (Ueno et al., 2011), HvHMA2 from barley (Mills et al., 2012), and CsHMA3/4 from cucumber (Migocka et al., 2015). The GxCCxxE motif, which occurs in the N or C terminus of all plant P_{1B2}-ATPases, is generally thought to be

1 associated with a heavy-metal-binding domain (Williams and Mills, 2005; Mills et al.,
 2 2010). In this study, a GxCCxxE motif was located in the N terminus of FIHMA3,
 3 thereby implying the presence of a heavy-metal-associated domain in this region (Fig.
 4 1). Overexpression of *OsHMA3* in a yeast mutant has been shown to affect sensitivity
 5 to Cd^{2+} and the ability to transport Cd^{2+} into vacuoles, which indicates that *HMA3* is
 6 responsible for sequestration of Cd^{2+} into vacuoles (Ueno et al., 2010; Miyadate et al.,
 7 2011). The methodologies we employed to demonstrate that *FIHMA3* similarly
 8 encodes a $\text{P}_{1\text{B}2}$ -ATPases transporter involved in sequestration of Cd^{2+} into the vacuole.

9 First, we fused GFP to FIHMA3 to visualize the subcellular localization of the
 10 protein in wild-type *A. thaliana* cells. Confocal imaging revealed that FIHMA3 is
 11 specifically located on the vacuolar membrane (Fig. 2). Previous studies have found
 12 that AtHMA3 and OsHMA3 are localized on vacuolar membranes and are involved in
 13 transporting heavy metal ions from the cytoplasm into vacuoles (Gravot et al., 2004;
 14 Ueno et al., 2010). The consistency observed in localization of FIHMA3 to vacuolar
 15 membranes suggests a possible role for the protein in the transfer of Cd^{2+} from the
 16 cytoplasm into the vacuole across the vacuolar membrane. Furthermore, gene
 17 expression analysis indicated that *FIHMA3* was mainly expressed in roots of *F.*
 18 *loliaceum* exposed to CdCl_2 stress (Fig. 3) as in the case of rice *OsHMA3* (Miyadate
 19 et al., 2011). An equivalent functional role may also be anticipated for *FIHMA3* as a
 20 determinant factor in the root in Cd tolerance. *FIHMA3* expression patterns in *F.*
 21 *loliaceum* do reflect a possible adaptation response to CdCl_2 stress. Ueno et al. (2010)
 22 reported that OsHMA3 from a low Cd^{2+} accumulating cultivar (Nipponbare) functions

as a firewall by sequestering Cd^{2+} into the vacuoles of roots, thereby separating Cd^{2+} from areal parts. A subsequent study confirmed that *OsHMA3* overexpression enhances tolerance to Cd^{2+} toxicity by increasing sequestration of Cd^{2+} into the vacuoles of root cells and then decreasing the translocation of toxic Cd^{2+} into shoots (Sasaki et al., 2014). One likely explanation is that only a limited amount of Cd^{2+} is loaded into the xylem from the root cells and subsequently translocated to the shoots (Miyadate et al., 2011). We note that a significant positive correlation was found between *FlHMA3* expression levels and Cd accumulation in roots of *F. loliaceum* exposed to different concentrations of Cd (10–100 mg L^{-1}) for 168 h or to 25 or 100 mg L^{-1} over the same time period (Figs. 4B–6B). This speculates that *FlHMA3* was detoxification of Cd^{2+} in *F. loliaceum* cells by enhancing the sequestration of Cd^{2+} into the vacuole.

Both shoots and roots of *F. loliaceum* plants reached their highest Cd contents at the highest applied CdCl_2 concentration (100 mg L^{-1}). The effect of this high concentration was morphologically reflected by a significant reduction in plant growth compared with the other Cd treatments (Figs. S2 and 4A). Most heavy-metal-accumulating plants generally exhibit slow growth rates, low biomass accumulation, and a tendency for altered root morphologies and necrosis (Clemens, 2006). Although the use of *F. loliaceum* for bioremediation of polluted soils requires further scrutiny, the pilot study described herein suggests that this grass hybrid has potential in such an application. One limitation of the current study is that our Cd treatments, although extremely concentrated and highly toxic to plant growth, were

only applied over 168 h. Future research using the same *F. loliaceum* cultivar should include much longer exposures to Cd stress. Assuming the promising results reported here are observed over a longer time period, this grass may be used simultaneously for two functions: bioremediation of Cd-contaminated land and provision of fodder for livestock (its original agricultural role). In the latter case, the uptake of Cd²⁺ into the shoots would have to be negligible to prevent harm to animals or subsequent entry into the food chain. An additional consideration is that other heavy metals are likely to be present in Cd-contaminated soils; the impact of these on *F. loliaceum* has yet to be assessed.

5. Conclusion

Our results demonstrate that *FIHMA3* encoding a P_{1B2}-ATPase is a tonoplast transporter that may play important roles in the response of *F. loliaceum* to Cd stress. Further research should focus on understanding the mechanism in detail by using methods like mutation or gene silencing.

Acknowledgements

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- 22

1 **Figure Legends**

2 **Fig. 1.** Sequence alignment of FIHMA3 with other HMA3s from higher plants.

3 Sources of P_{1B2}-ATPases and their GenBank accession numbers are as follows:

4 AtHMA3 (*Arabidopsis thaliana*, NM_119158), OsHMA3 (*Oryza sativa*,

5 XM_015791882), and TaHMA3 (*Triticum aestivum*, KF683298). The sequences were

6 aligned using DNAMAN 8.0 software. Amino acid residues highlighted in black are

7 conserved in the three transporters. Identical and different amino acid residues are

8 indicated with white and blue, respectively. Eight putative transmembrane domains

9 (TM 1–TM 8) and several motifs are boxed.

10 **Fig. 2.** Subcellular localization of an FIHMA3-GFP fusion protein transiently

11 expressed in *Arabidopsis thaliana* mesophyll cells. (A) Images obtained using GFP

12 alone as the control. The red dye FM 4-64 was used to indicate the plasma membrane

13 location. (B) Images obtained when GFP was fused to the C terminus of FIHMA3. In

14 panels A and B from left to right, GFP signals, FM 4-64 signals, merged images of

15 GFP and FM 4-64 signals, and bright-field differential interference contrast (DIC)

16 images are shown. Bar = 5 μ m.

17 **Fig. 3.** *FIHMA3* expression in *F. liliaceum* exposed to (A) different concentrations of

18 CdCl₂ (0, 10, 25, 50, and 100 mg L⁻¹) for 168 h or to 25 mg L⁻¹ CdCl₂ (B) or 100 mg

19 L⁻¹ CdCl₂ (C) over a 168-h period. The relative expression level of *FIHMA3* in shoots

20 and roots was analyzed by quantitative real-time RT-PCR. *Actin* was used as an

21 internal control. Experiments were repeated three times. Data are means \pm SD ($n = 3$)

22 and bars indicate SD. Different letters indicate significant differences at $P < 0.05$

23 (Duncan's test).

Fig. 4. Cd content of tissues of *F. loliaceum* exposed to 10, 25, 50, and 100 mg L⁻¹ CdCl₂ for 168 h. (A) Cd content of shoots and roots. Five plants were pooled per replicate ($n = 8$). (B) Relationship between relative *FlHMA3* expression and Cd content of roots subjected to 10 (◆), 25 (▲), 50 (■), and 100 (●) mg L⁻¹ CdCl₂ treatment for 168 h ($n = 3-8$). Data are means \pm SD and bars indicate SD. Different letters indicate significant differences at $P < 0.05$ (Duncan's test).

Fig. 5. Time course of Cd content of *F. loliaceum* exposed to 25 mg L⁻¹ CdCl₂ for 3 to 168 h. (A) Cd content of shoots and roots. Five plants were pooled per replicate ($n = 8$). (B) Relationship between relative *FlHMA3* expression and Cd content of roots subjected to 25 mg L⁻¹ CdCl₂ treatment for 3–168 h. Data are means \pm SD ($n = 3-8$) and bars indicate SD.

Fig. 6. Time courses of Cd content of *F. loliaceum* exposed to 100 mg L⁻¹ CdCl₂ for 3 to 168 h. (A) Cd content of shoots and roots. Five plants were pooled per replicate ($n = 8$). (B) Relationship between relative *FlHMA3* expression and Cd content of roots subjected to 100 mg L⁻¹ CdCl₂ treatment for 3–168 h. Data are means \pm SD ($n = 3-8$) and bars indicate SD.

Supplementary Figure Legends

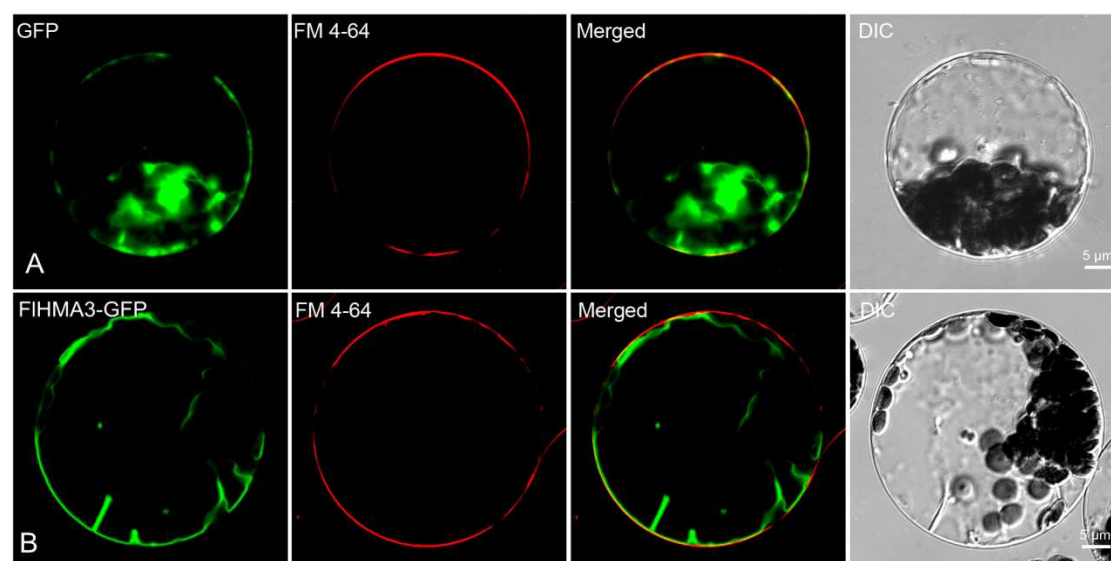
Supplementary Fig. 1. Phylogenetic tree of *HMA*s. The tree was constructed by the neighbor-joining method. Genes and GenBank accession numbers are as follows:

AdHMA3 (*Arachis duranensis*, XM_016078710), *AiHNA3* (*Arachis ipaensis*, XM_016346129), *AhHMA3* (*Arabidopsis halleri*, AJ556182), *AhHMA4* (*Arabidopsis halleri*, AY960757), *AtHMA2* (*Arabidopsis thaliana*, NM_119157), *AtHMA3*

1 (*Arabidopsis thaliana*, NM_119158), *AtHMA4* (*Arabidopsis thaliana*, AF412407),
 2 *BdHMA3* (*Brachypodium distachyon*, XM_003561234), *BnHMA3* (*Brassica napus*,
 3 XM_013849300), *BrHMA3* (*Brassica rapa*, XM_009139644), *CaHMA3* (*Cicer*
 4 *arietinum*, XM_012717947), *CsHMA3* (*Camelina sativa*, JX402100), *CmHMA3*
 5 (*Cucumis melo*, XM_008455480), *EgHMA3* (*Elaeis guineensis*, XM_010928912),
 6 *EugHMA3* (*Eucalyptus grandis*, XM_010048654), *ErgHMA3* (*Erythranthe guttata*,
 7 XM_012995791), *FlHMA3* (*Festulolium loliaceum*), *FvHMA3* (*Fragaria vesca*,
 8 XM_011464053), *GmHMA3* (*Glycine max*, XM_006593460), *GrHMA3* (*Gossypium*
 9 *raimondii*, XM_012589041), *HvHMA2* (*Hordeum vulgare*, GU177852), *HvHMA3*
 10 (*Hordeum vulgare*, KU212808), *JcHMA3* (*Jatropha curcas*, XM_012211439),
 11 *MaHMA3* (*Musa acuminata*, XM_009417251), *MnHM3* (*Morus notabilis*,
 12 XM_010112413), *NbHMA3* (*Nicotiana tabacum*, XM_016654239), *NcHMA4*
 13 (*Noccaea caerulescens*, JQ904704), *ObHMA3* (*Oryza brachyantha*, XM_006658354),
 14 *OsHMA2* (*O. sativa*, HQ646362), *OsHMA3* (*O. sativa*, XM_015791882), *PdHMA3*
 15 (*Phoenix dactylifera*, XM_008803179), *PeHMA3* (*Populus euphratica*,
 16 XM_011021683), *PmHMA3* (*Prunus mume*, XM_008225567), *RcHMA3* (*Ricinus*
 17 *communis*, XM_015727254), *SaHMA2* (*Sedum alfredii*, JQ012929), *SbHMA3*
 18 (*Sorghum bicolor*, XM_002459533), *SiHMA3* (*Setaria italica*, XM_012843843),
 19 *SlHMA3* (*Solanum lycopersicum*, XM_004242795), *TaHMA2* (*Triticum aestivum*,
 20 HM021132), *TaHMA3* (*Triticum aestivum*, KF683298), *ThHMA3* (*Tarenaya*
 21 *hassleriana*, XM_010550291), *TtHMA3* (*Triticum turgidum*, KF683295), *VvHMA3*
 22 (*Vitis vinifera*, XM_010658478), and *ZmHMA3* (*Zea mays*, XM_008671782).

- 1 **Supplementary Fig. 2.** Relative growth rate (RGR) of *F. loliaceum* exposed to 0, 10,
2 25, 50, or 100 mg L⁻¹ CdCl₂ for 168 h. Five plants were pooled per replicate ($n = 8$).
3 Data are means \pm SD and bars indicate SD. Different letters indicate significant
4 differences at $P < 0.05$ (Duncan's test).

31

**Fig. 2**

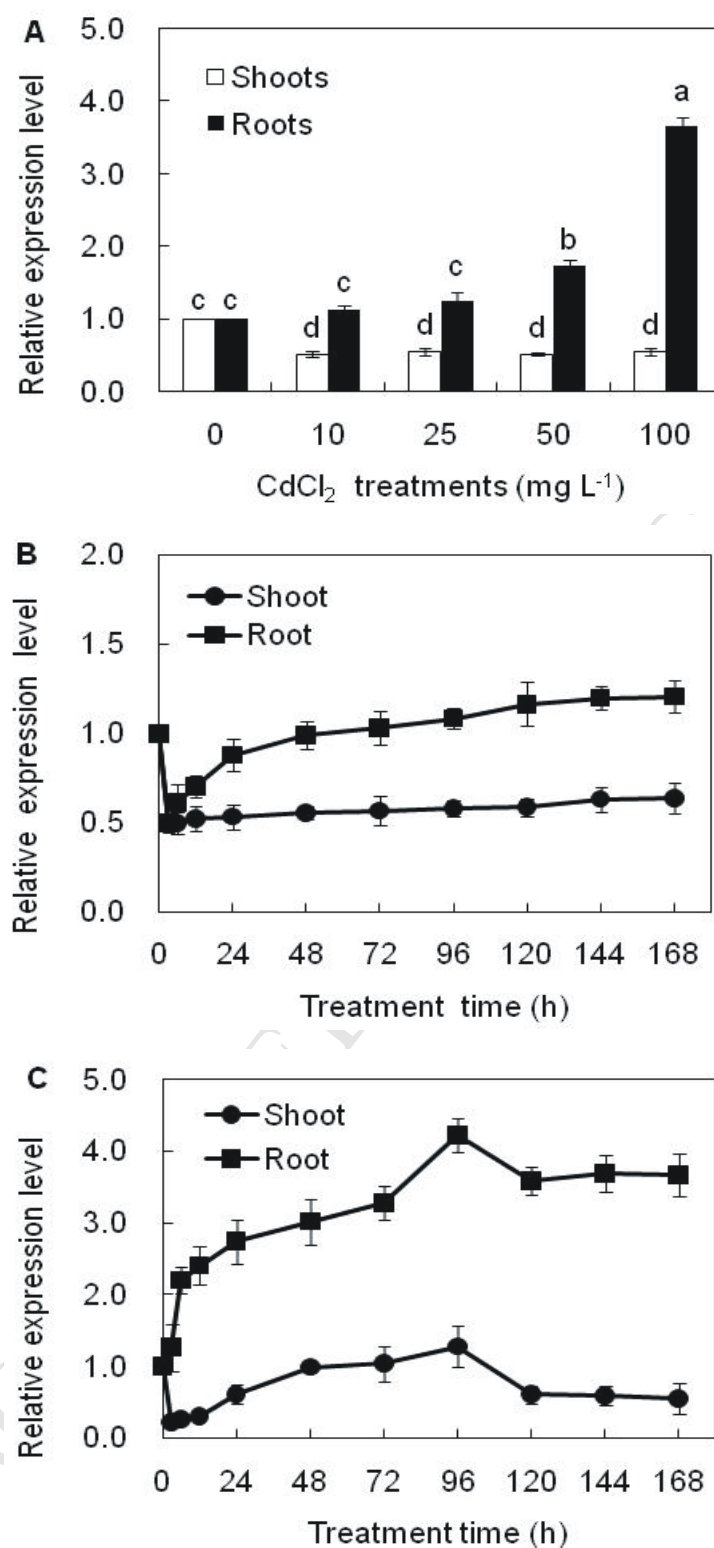
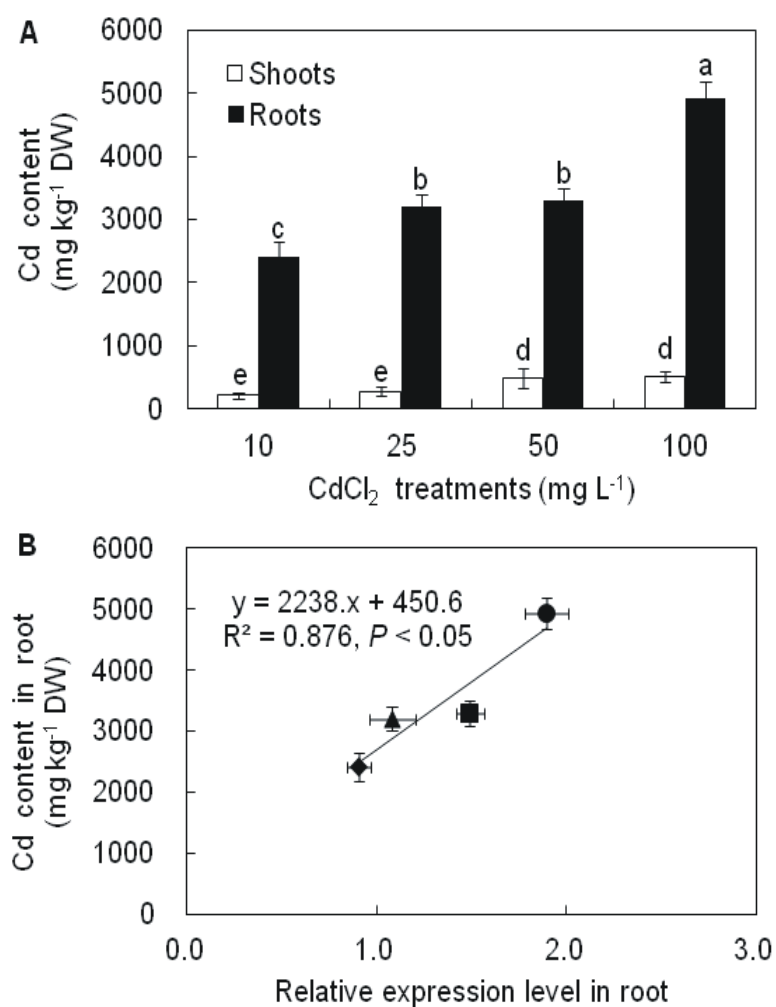


Fig. 3

**Fig. 4**

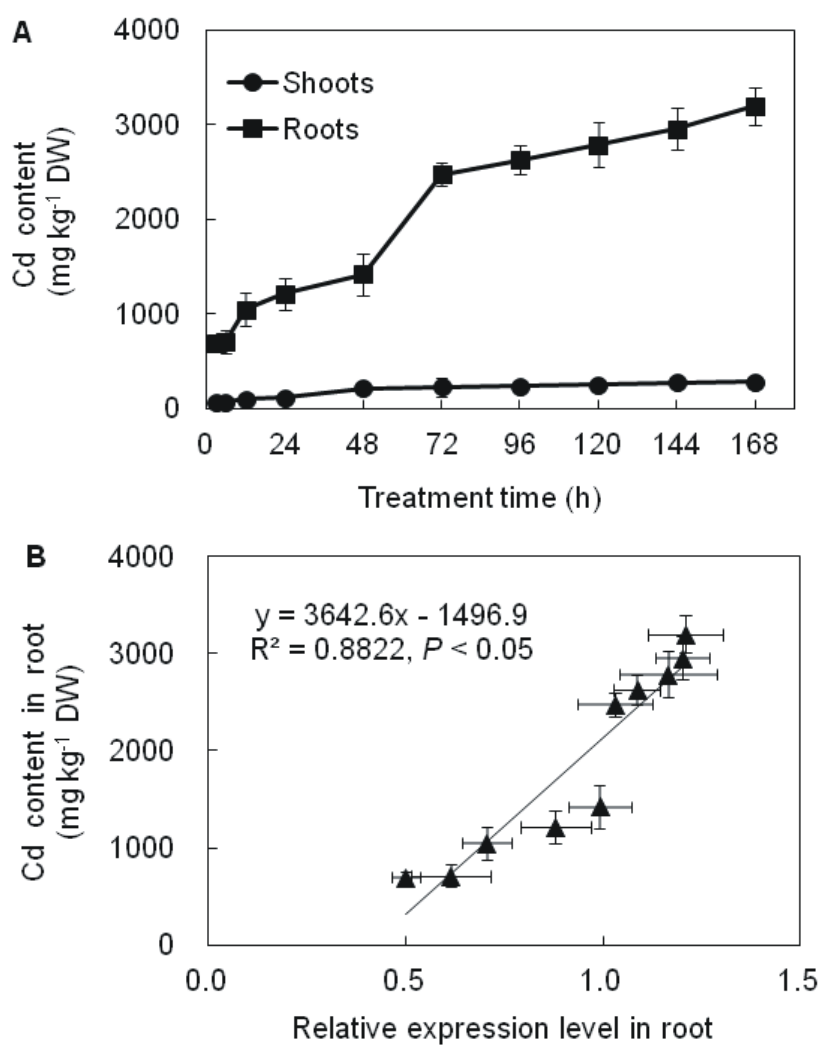
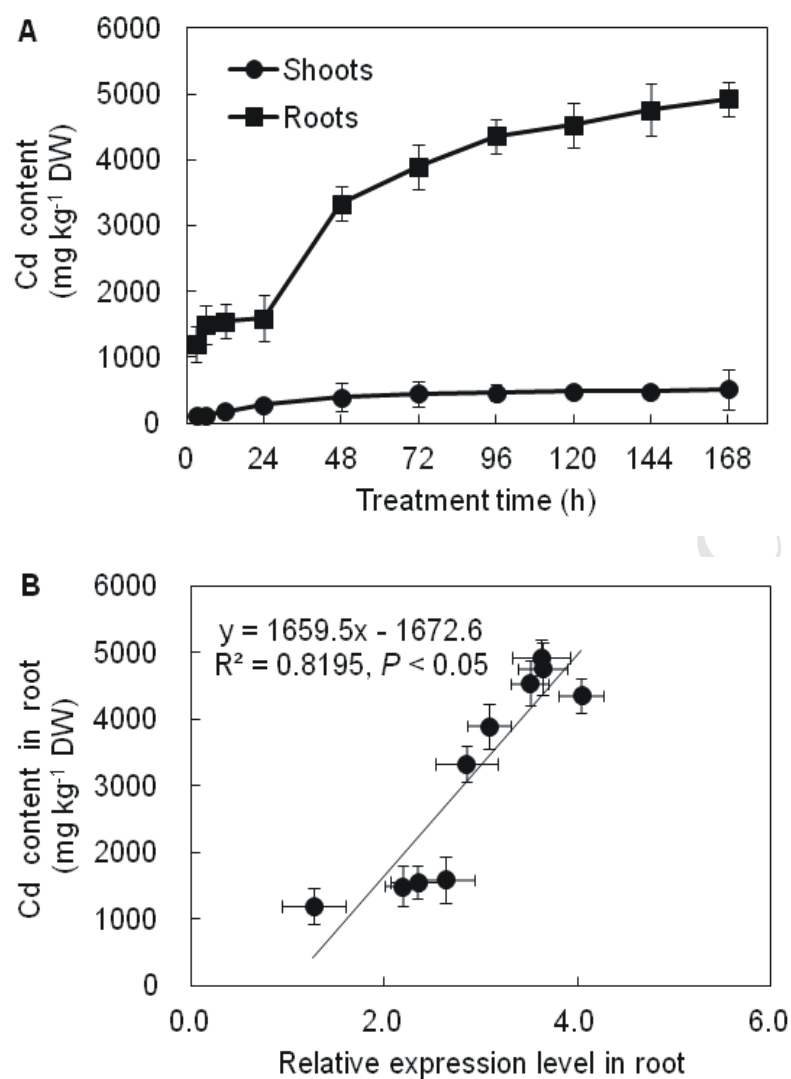
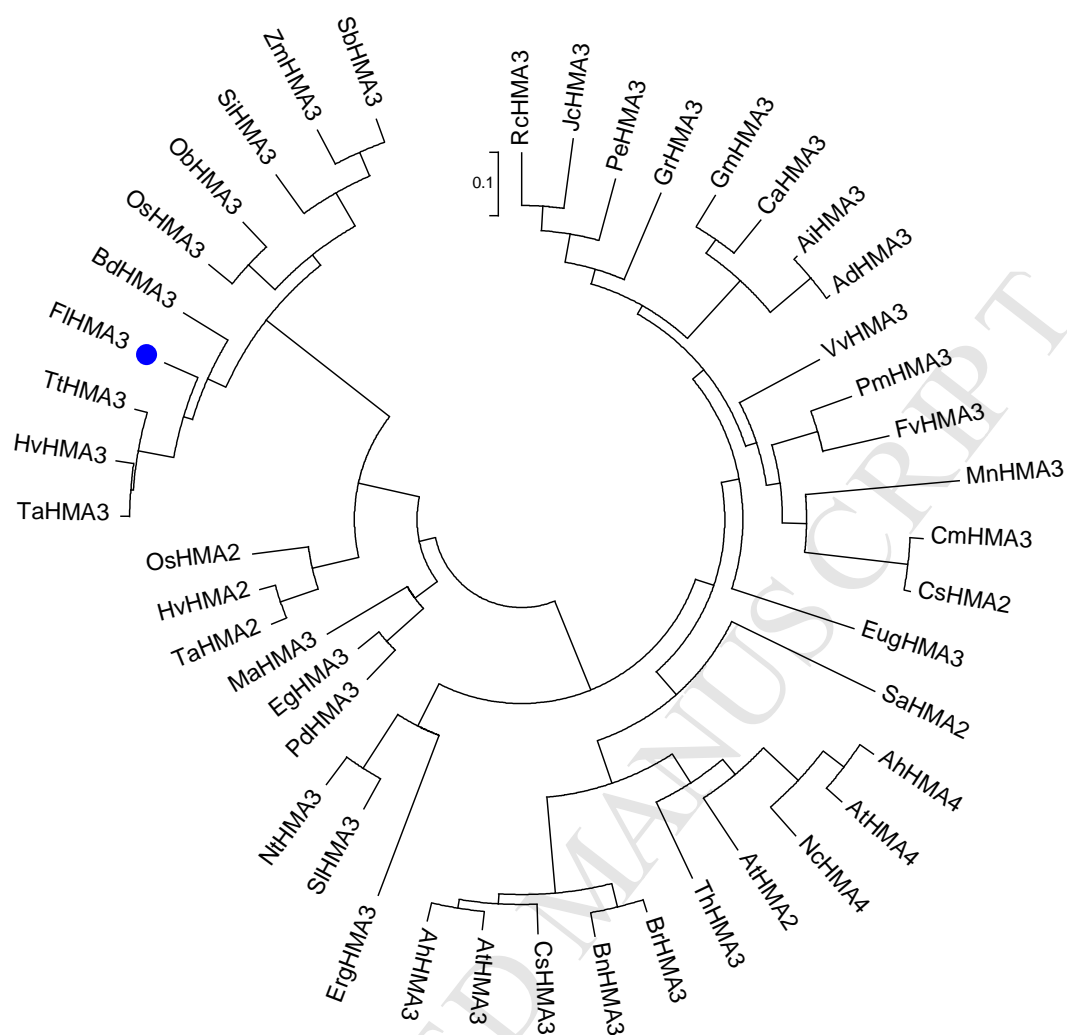
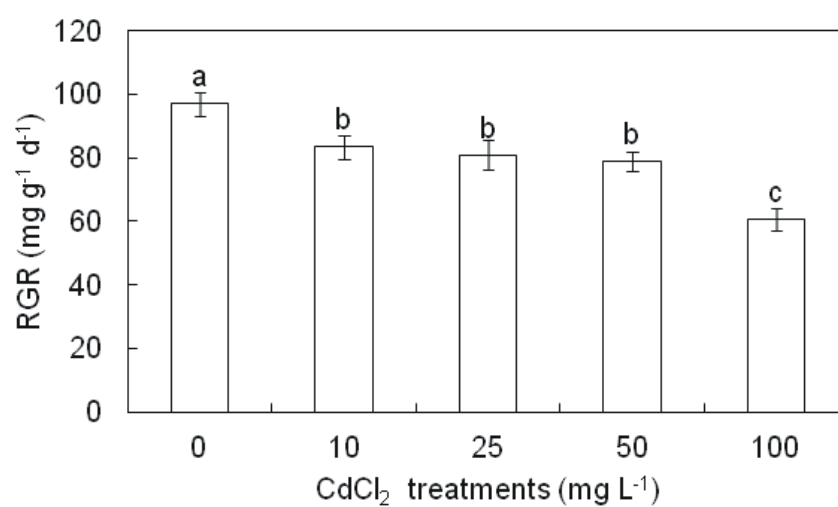


Fig. 5

**Fig. 6**



Supplementary Fig. 1

**Supplementary Fig. 2**

Supplementary Table 1

Primer sequences used in the experiments

Primer	Sequence (5'-3')
P1	ATCAACRTYCTSATGCTYATCGC
P2	GGTGATSGTSCCGGTCTTGTCGA
P3	GGTTACATTGCCGTGAGGACGAC
P4	TCGACAAGACCGGCACCATCACC
P5	GGGGAATTCATGACGGACGGTGGCGAGAAC
P6	GGGGGTACCGTTTCACCAGAGCAGCATCCT
P7	GCTCAACCTGGACGGTTACA
P8	AACCATCGCTCCAGATCACC
A1	TCGAGACTGCGAAGAGTAGC
A2	TCCATGCCGATGATGGAAGG

Highlights

- *FIHMA3* was mainly expressed in roots and up-regulated by excess Cd.
- FIHMA3 was localized at the vacuolar membrane.
- A significant positive correlation was found between expression levels of *FIHMA3* and Cd accumulations in roots of *F. loliaceum* under Cd stress.
- Cd²⁺ taken up by root cells may be sequestered into the vacuole via a pathway mediated by FIHMA3 to reduce the concentration of this toxic metal in the cytoplasm.

Author contributions

Qiang Guo, Lin Meng and Mike W. Humphreys conceived and designed the experiments. Qiang Guo performed all the experiments and wrote the manuscript. John Scullion and Luis A.J. Mur review and polish the manuscript.